



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of
A. Dömling

Application No.: 10/520,791

Filed: January 8, 2005 Art Unit: 1654

For: TUBULYSIN CONJUGATES Examiner: S.R. Gudibande

Commissioner for Patents
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DECLARATION UNDER 37 CFR 1.132

I, Alexander Dömling, declare as follows:

1. I am the Inventor on the above-identified patent application (referred to below as "the patent application"). I earned a Ph.D. degree in Chemistry from the Technical University in Munich in 1993. Subsequently, I was Vice-President Chemistry of Morphochem AG until 2003 and then in 2004 co-founded R&D Biopharmaceuticals GmbH. I am currently Associate Professor of Pharmacology at the University of Pittsburgh.

2. The following experimental work as detailed in paragraph 3 below and in the enclosed poster hand-out were conducted by me or persons working under my direction.

3. The tubulysin compounds identified in the table below were tested in an acid phosphatase assay for activity against human cancer cell lines of MCF-7 and KB-V1. The protocol of the acid phosphatase assay was as described in Yang et al., Anal. Biochem. 241 (1996) 103.

No	Structure	Code	MCF-7 IC50 [ng/ml]	KB-V1 IC50 [ng/ml.]
1		Tubulysin A	0.7	1.0
2		Tubulysin D	0.5	0.3

4. I declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application of any patent issuing therein.

Date: 30/06/07

Alexander Dömling

Enclos.:

- poster hand-out "Preclinical antitumor activity of Polymer, Tubulysin Nanoparticles in Human Colorectal Cancer Xenograft"

Preliminary antitumor activity of Polymer-Tubulysin Nanoparticles in Human Colorectal Cancer Xenograft

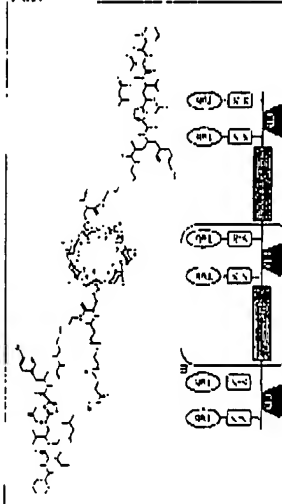
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I. Introduction

Tub A derivative-Tubulysin nanoparticles are conjugates of a Tubulinysin A (Tub A) derivative and a linear, cycloheximide-based peptide [C22]. A similar polymer-cycloheximide conjugate is in clinical trials now for cancer treatment. Tub A is a naturally occurring helicopeptide isolated from strains of mycobacteria. It is highly active against multiple cancer cell lines with an IC₅₀ in the low nM to pM concentration range. It acts as an antimicrotubule agent that depolymerizes cell microtubules and triggers apoptosis. TR = *L*-A derivative's covalently attached to CDP through a disulfide linker (Figure 1).

Figure 1. Structure of CDP-S-S-Tub



II. Characterization and Release Studies

The Tub A derivative was incorporated to the polymer backbone with a loading of 12% by weight as measured by HPLC. The particle size of the parent polymer was measured to be 10 nm while CDP-S-S-Tub self-assembled into nanoparticles with a particle size of 12 nm. The solubility of Tub A in water was determined to be 0.1 mg/ml at a neutral pH while that of CDP-S-S-Tub was found to be 100 times higher.

Release studies was performed by incubating CDP-S-S-Tub in both PBS and human plasma. The conjugate was found to be stable at both conditions for greater than 72 h at 37°C.

III. In vitro Studies

The anti-proliferative activity of CDP-S-S-Tub was evaluated in vitro in multiple human cancer cell lines (Table 1). The data shows that the conjugate maintains high anti-proliferative activity.

Table 1: IC₅₀ studies

Cell Type	IC ₅₀ (nM)	Tub A	CDP-S-S-Tub
NCI-H1229 (lung) cells	2.3	2.8	4.4
HT-29 (colon) cells	4.9	1.3	4.4
A2780 (ovarian) cells	2.5	2.4	N/A

IV. MTD Studies

The maximum tolerated dose (MTD) of CDP-S-S-Tub was determined in nude mice and found to be between 3 to 10 mg/kg (in Tub equivalents) whereas that of Tub A was not yet established due to 100% mortality at a dose of 0.3 mg/kg (Table 2).

Table 2: MTD studies

Group	n	Treatment agent (mg/kg)	Mean BW (g)	% of TR	Avg Day of TR
1	4	CDP-S-S-Tub	10	[23.4%]	5
2	4	Tub A	3	[9.7%]	1
3	4	CDP-S-S-Tub	1	[0.4%]	Day 2
4	4	Tub A	3	[8.7%]	Day 2
5	4	Tub A	1	[1.7%]	Day 3
6	4	Tub A	0.3	[0.1%]	Day 8

* All mice were treated with subcutaneous injection of 100 µg/kg.

** TR = Indicated relative toxicity.

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V. Efficacy Studies

Preliminary efficacy was evaluated in nude mice bearing subcutaneously implanted HT-29 colorectal xenografts. Treatment with CDP-S-S-Tub was well-tolerated with no mortality and significant antitumor effect. It was believed that the conjugate showed high antiproliferative activity in nude mice compared to Vinblastine. Treatment with Tub A was proven to be toxic to the mice, causing 50% mortality and 28.8% max tumor body weight loss on day 26 (Table 3 and Graph 1 & 2).

Table 3: Summary of antitumor activity (End point TV = 1000 mm³ or Day 26, whichever comes first)

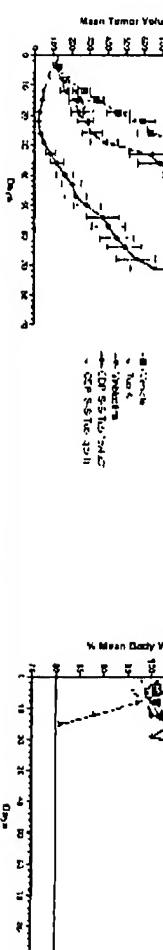
Group	n	Agent	Drug Schedule	MTD (mg/kg)	BW (g)	Median TTE	TC	MTG3	Statistical Significance ^a		Regressions: Deaths
									vs G1	vs G4	
1	10	Vehicle	—	CWx3	—	—	33.65	—	—	—	0
2	10	Tub A	D1	CWx3	—	—	25.82	34.46	0.8*	2.42	0
3	10	Vinblastine	4	QWx3	—	—	3.60%	45.05	11.4	33.80	0
4	10	CDP-S-S-Tub	3 ^b	QWx3	700.0 (3)	—	2.20%	73.55	11.87	***	0
5	10	CDP-S-S-Tub	3 ^c	QWx3	—	—	2.90%	56.55	23.27	11.91	—

^a All p-values were generated using Kaplan-Meier (Graph 1), Tukey HSD (30% survival) or Kruskal-Wallis (Graph 2) for the number of animals surviving by endpoint. ^b Control group received 10% DMSO v/v. Tukey HSD 30% survival p-value = 0.0000000000000001. ^c Tukey HSD 30% survival p-value = 0.0000000000000001.

Graph 1: Tumor Growth Daily in HT-29 xenograft implanted mice



Graph 2: % Mean Body Weight Loss over time in HT-29 xenograft implanted mice



VI. Conclusion

A polymer-disulfide conjugate CDP-S-S-Tub was synthesized and found to be highly soluble in water and stable in both PBS and human plasma.

The conjugate showed high antiproliferative activity in null mice bearing human cancer cell lines.

Efficacy studies of CDP-S-S-Tub at 3 months showed that it was well-tolerated and provided substantial therapeutic activity during a 90 day study. By contrast, the free drug Tub A, showed excessive toxicity.

Vinblastine, a vincristine alkaloid that inhibits tubulin polymerization by binding to the same binding site as Tub A was significantly less effective as an antitumor agent compared to CDP-S-S-Tub.

VII. Acknowledgments

We thank Piedmont Research Center for performing all of the animal studies.

PA Boeters & Lieck